

Bioinformatics for high-throughput data

Today's Outline

Part A : Introduction on high-throughput data
Experimental design
Platforms for gene expression data

Break – 10 min (questions if any)

Part B : Exponential and logs
Low level analysis

- Gene Expression Estimation
- Normalisation

Class activity: Research on burning questions.

Today we will be learning:

- Characteristics of high-throughput data and how it is produced
- What we intend for OMICS
- How we define high-throughput data
- To define principles of experimental design and pipelines
- Methods for gene expression quantification
- To estimate gene expression levels from data and difference in methods applied
- To normalise the data and what it means

High-throughput data

High-throughput data is a large amount of data collected using automated methods and non-conventional technologies. This is to be able to perform a large number of experiments at the same time to monitor a behavior of a system.

High-throughput data is produced in all field where quantitative science is applied. For example for robotics and automated system, in ecology to study and monitor populations, in chemistry for compound screening and obviously in biology.

High throughput *cell biology* is the use of automation equipment with classical cell biology techniques. This is to address biological questions that are otherwise unattainable using conventional methods.

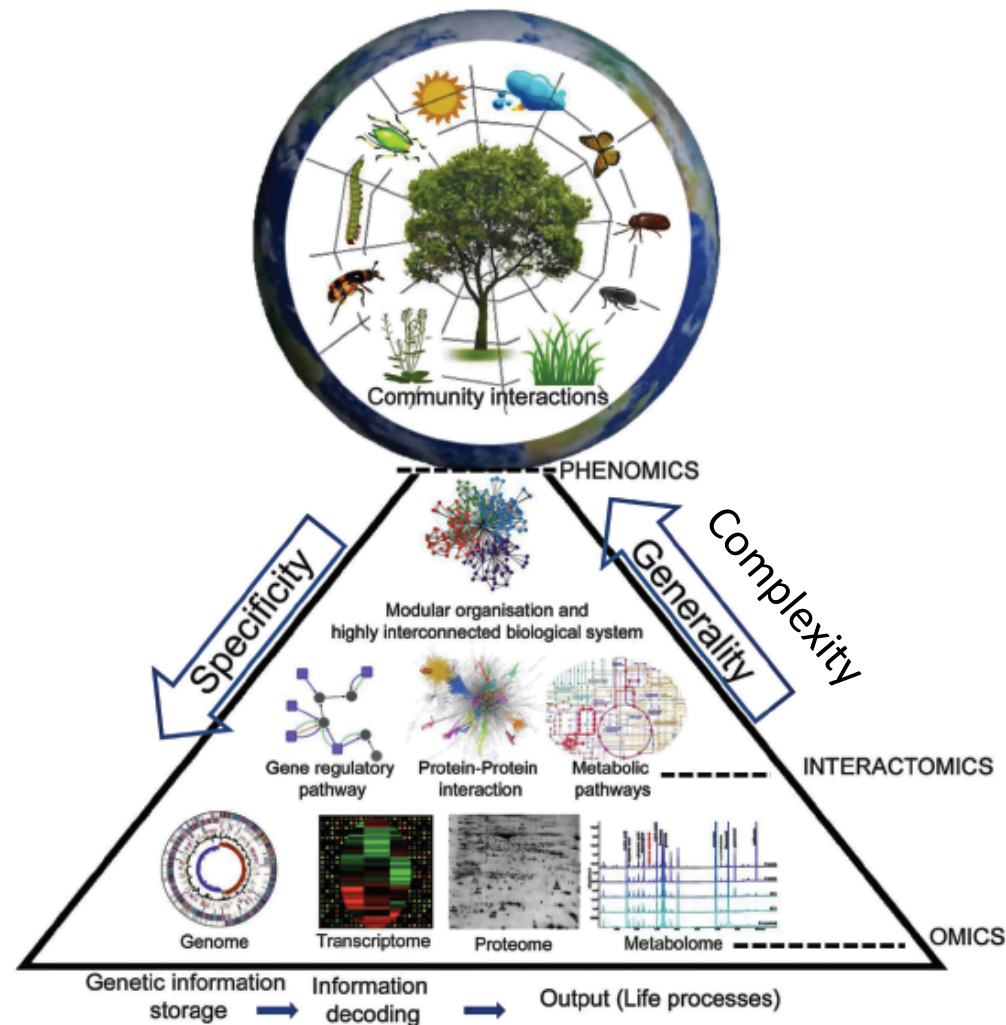
High-throughput biology has created a new field of biology called *OMICS*. It is a research field that interface between large scale biology (genome, proteome, transcriptome), technology and computational methods.

Characteristics of high-throughput data

What are the characteristics of high-throughput data?

- Large in size
- Prone to many false positives (low specificity)
- Capture biological Noise
- Accurate
- Technically noisy

Omics and its hierarchy



To microscopic to
macroscopic

The highly interconnected hierarchical organization and functional complexity need a sophisticated integrated systems approach.

Why do we need mathematics to understand biology?

Quantification of genome wide gene expression
Gene networks and target predictions
Protein-Protein Interaction Networks

Biological systems

Alignment tools
Mining of large data
Accessing large data from repositories

Large Data

Handling complexity
non-integer dimensions
Self-similarity --- Patterns

Patterns and structures

Quantification of Uncertainty: Be able to predict
what we cannot measure and have a theoretical
way of quantify accuracy

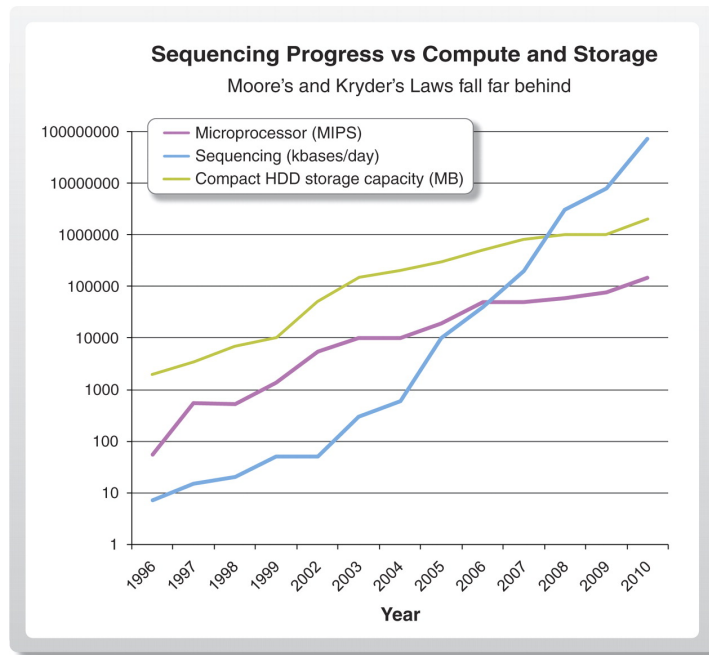
Modeling

Computational Biology --- Bioinformatics

Genomic Data: the truth about it

In February 2001 scientists published the first drafts of the Human Genome. The dawn of the genomic era.

Challenges in genomics derive mainly from the *informatics* that is required to **analyse, summarised** and **interpret** the vast amount of raw (sequencing and not) data that is available.



S D Kahn, Science 2011;331:728-729

Moore's law: The number of transistors that can be placed inexpensively on an integrated circuit doubles approximately every two years

Current information technology allows to perform standard analysis of raw output from sequencing in average time of 3 days on a cluster

Value and definition of raw data?
How can we analyse this data effectively?

Tools we need

- Optimal Experimental Design – minimise the noise in the measurement
- Mathematical Models – define rules (functions) to describe processes
- Statistical tools – quantify accuracy in prediction and sensitivity in estimation
- Computational Skills – handling large amount of data in automated way
- Visualisation tools – identify patterns in the data

The role of experimental design

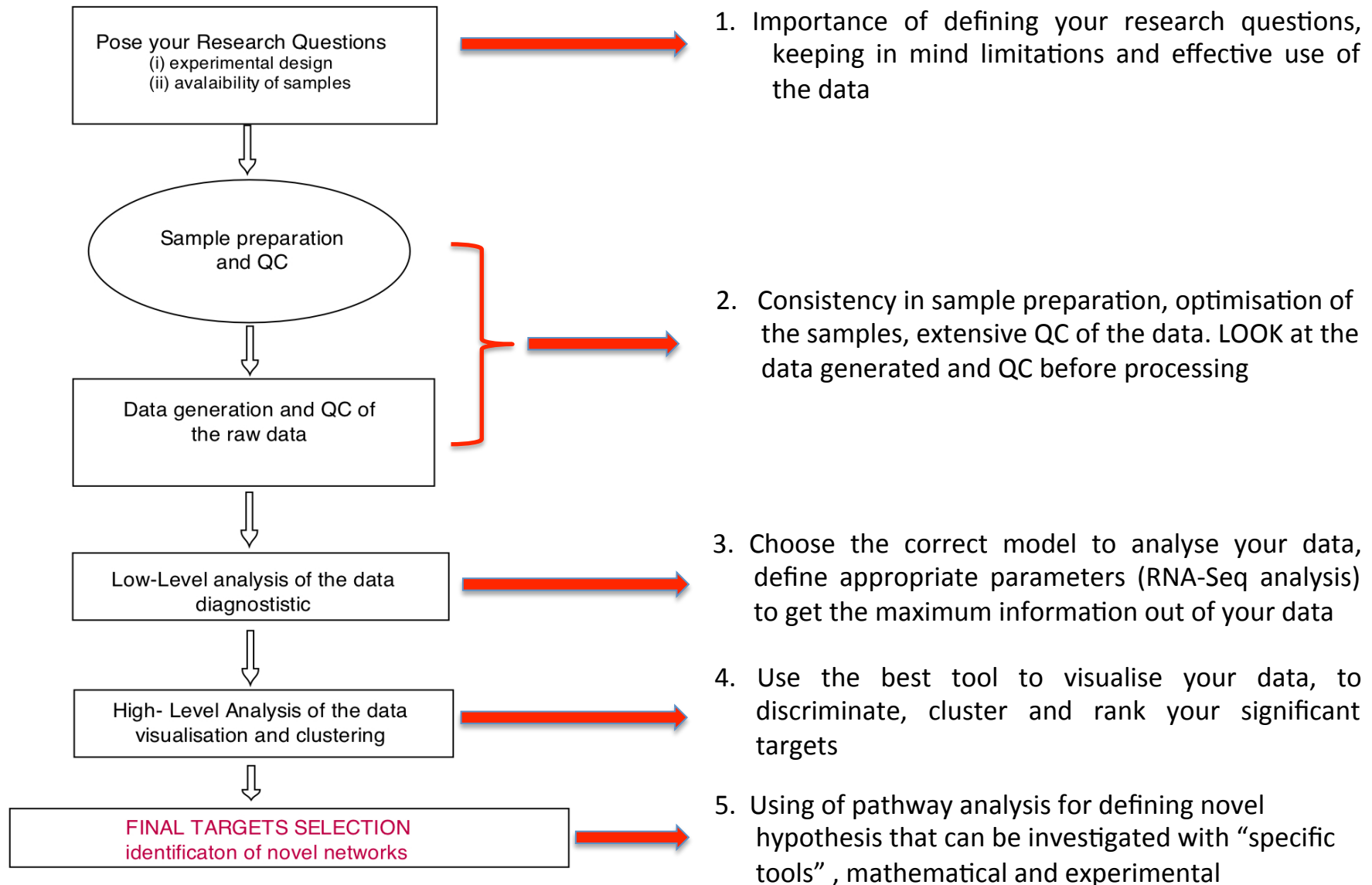
Importance of defining your research questions, keeping in mind limitation and effective use of the data

We need to limit the uncertainty of the “unknown” by define very clear questions. This helps to :

- reduce variability in the data.
- reducing the complexity of the data by focusing the search of information on the questions you have in mind.

A proper experimental design **MUST** reflect :

- the biological questions that you are asking,
- the protocols optimised to minimise the variations in the data
- identify the limitations of the data collected.



RNA sample preparation for high-throughput assays

Experimental design:

your RNA collection needs to reflect the biological questions you are asking

Optimise protocols:

- minimise technical errors
- minimise batch effects
- clean and pure sample to avoid contamination

Estimate the correct quantities:

- samples for validation
- optimisation of the protocols – avoid saturation or low quantification

Technical and biological replicates:

- ensure you have SOP in place

Pipelines

All the variation and to ensure that the analysis of the data is as reproducible as the experimental collection of samples generate the need to define pipelines for the analysis of the data

PIPELINES: Reproducible and robust protocols for numerical experimentations. In case of biological data they are tailored to the system/organism under study.

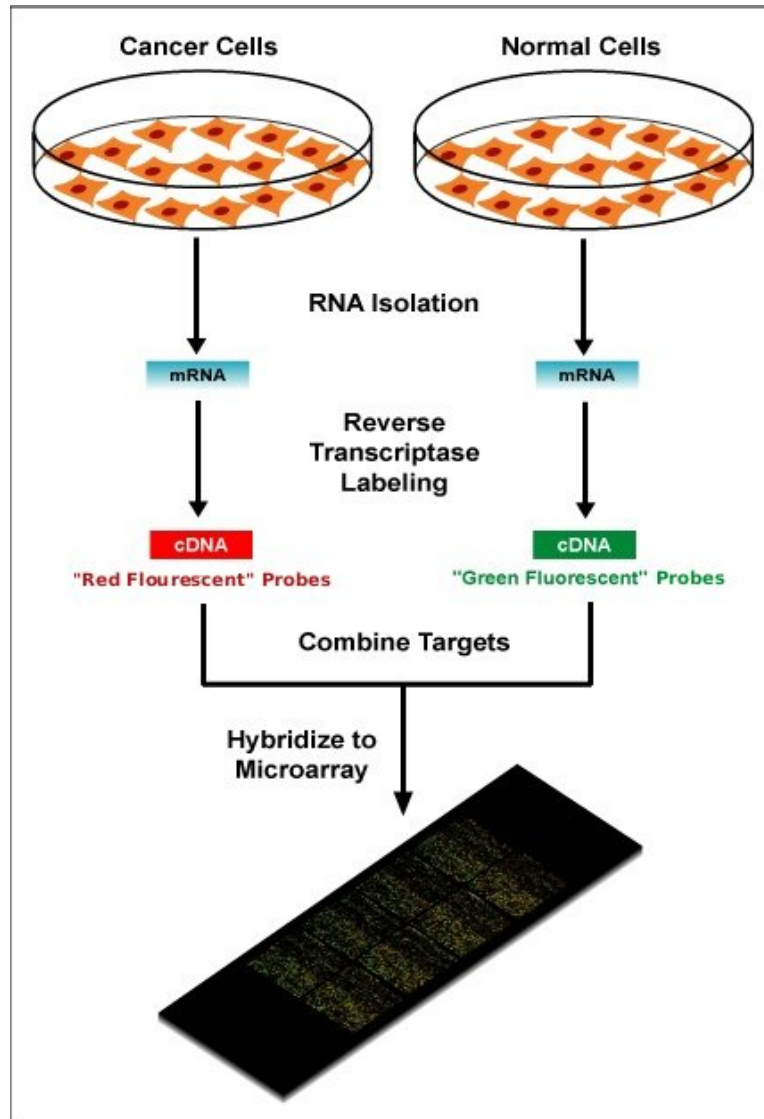
HOW DO WE GET THEM?

Gene expression estimation

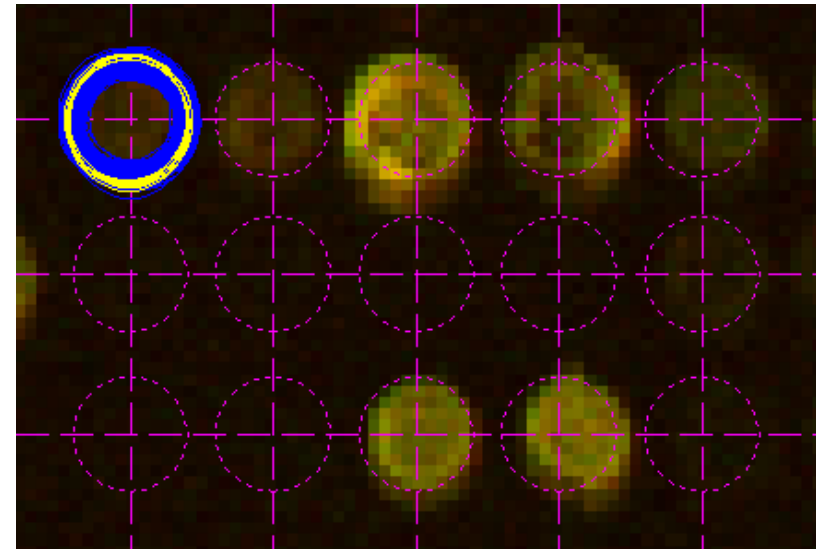
Interpret and analyse the data by first understanding where the data is coming from

Different platforms that generate gene expression:

- Two or One color spotted cDNA arrays
- Affymetrix - new Human/Mouse whole transcriptome arrays
- Illumina Arrays
- Reads – RNA-Seq and other NGS assays

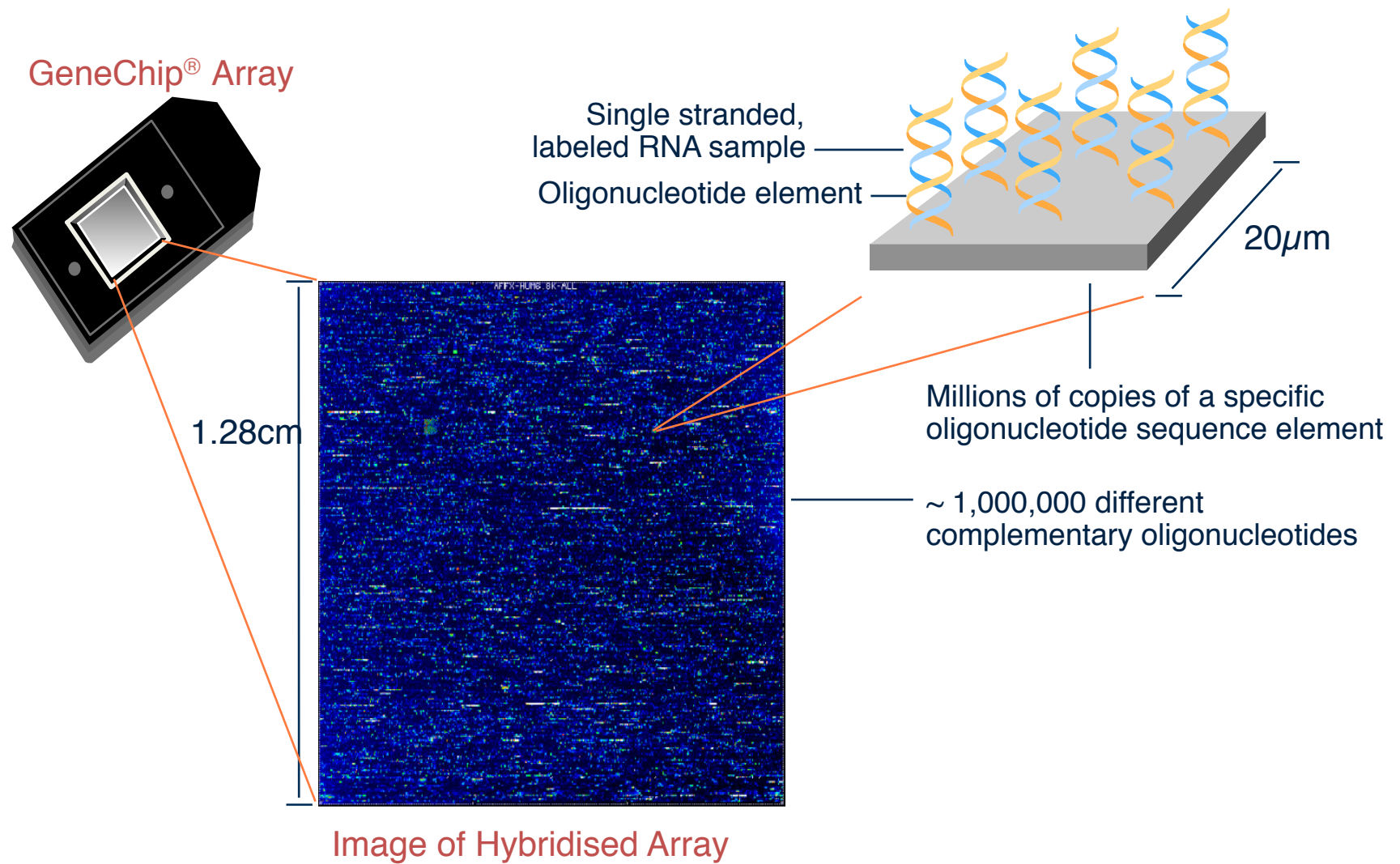


Wikipedia – DNA microarrays



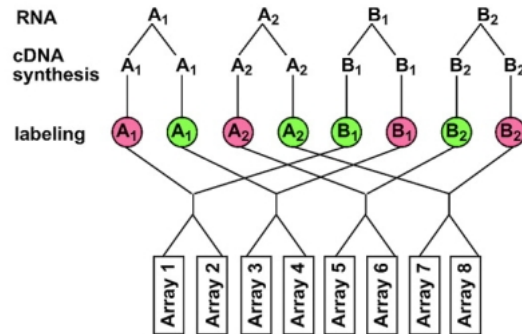
How do I quantify the gene expression?

Image processing noise
 Sensitivity

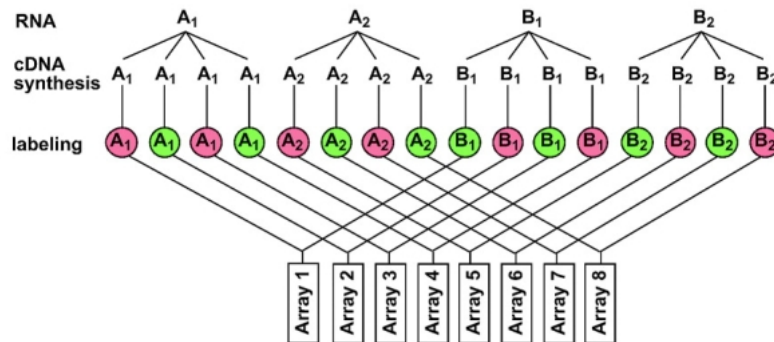


Example of experimental design

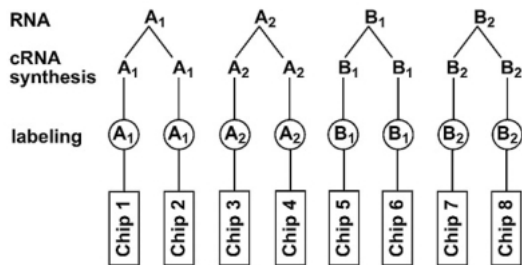
A. cDNA



B. Oligo



C. Affymetrix



- Relative expression
- Important to choose the reference
- Important to choose the experimental design

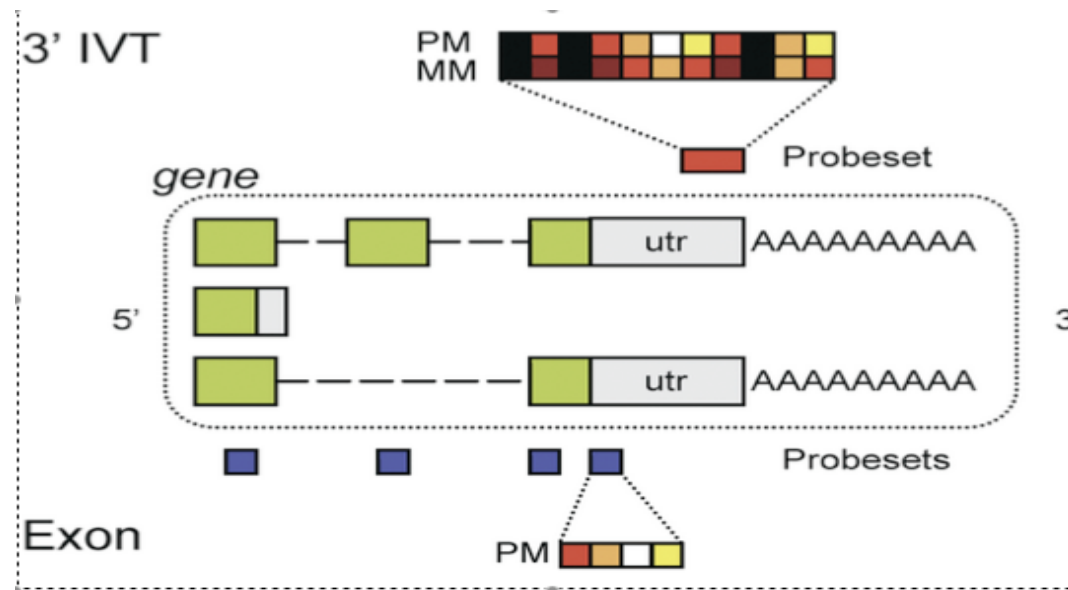
- Estimation of Absolute expression
- No reference
- Not specific to the question you are asking
- Important the design – data analysis

Example: HG_U133 Plus v2 Affymetrix geneChip

The sequences from which these probe sets were derived were selected from GenBank®, dbEST, and RefSeq.

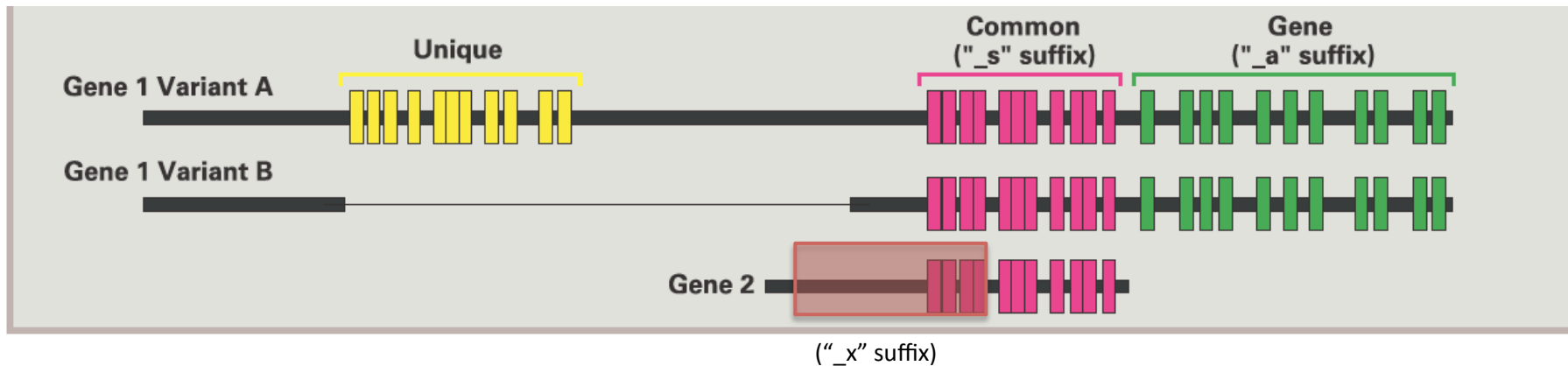
A single array contains with more than 54,000 probe sets representing approximately 38,500 genes (estimated by UniGene coverage).

70 percent of the probe sets represent subcluster assemblies containing one or more non- EST sequences. Of the 16,737 EST-based probe sets, approximately 9,000 probe sets can now be associated with an mRNA or other non-EST sequence.



Now with new arrays HJAY.....

Probe Set Notation



_at : probe sets are predicted to perfectly match only a single transcript

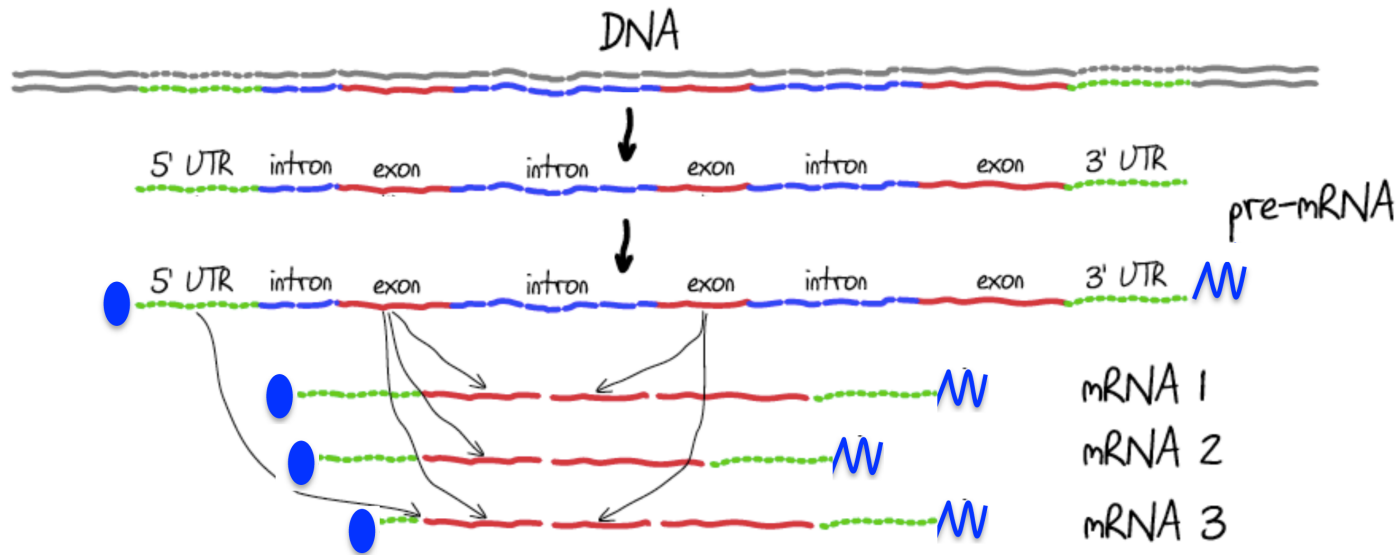
_s_at : are predicted to perfectly match multiple transcripts, which may be from different genes

a_at : all probes in the probe set hit alternate transcript of the same gene

_x_at : probe sets will contain some probes that are identical or highly similar to other sequences from different gene.

Hybridize uniformly across probe pairs to the intended target

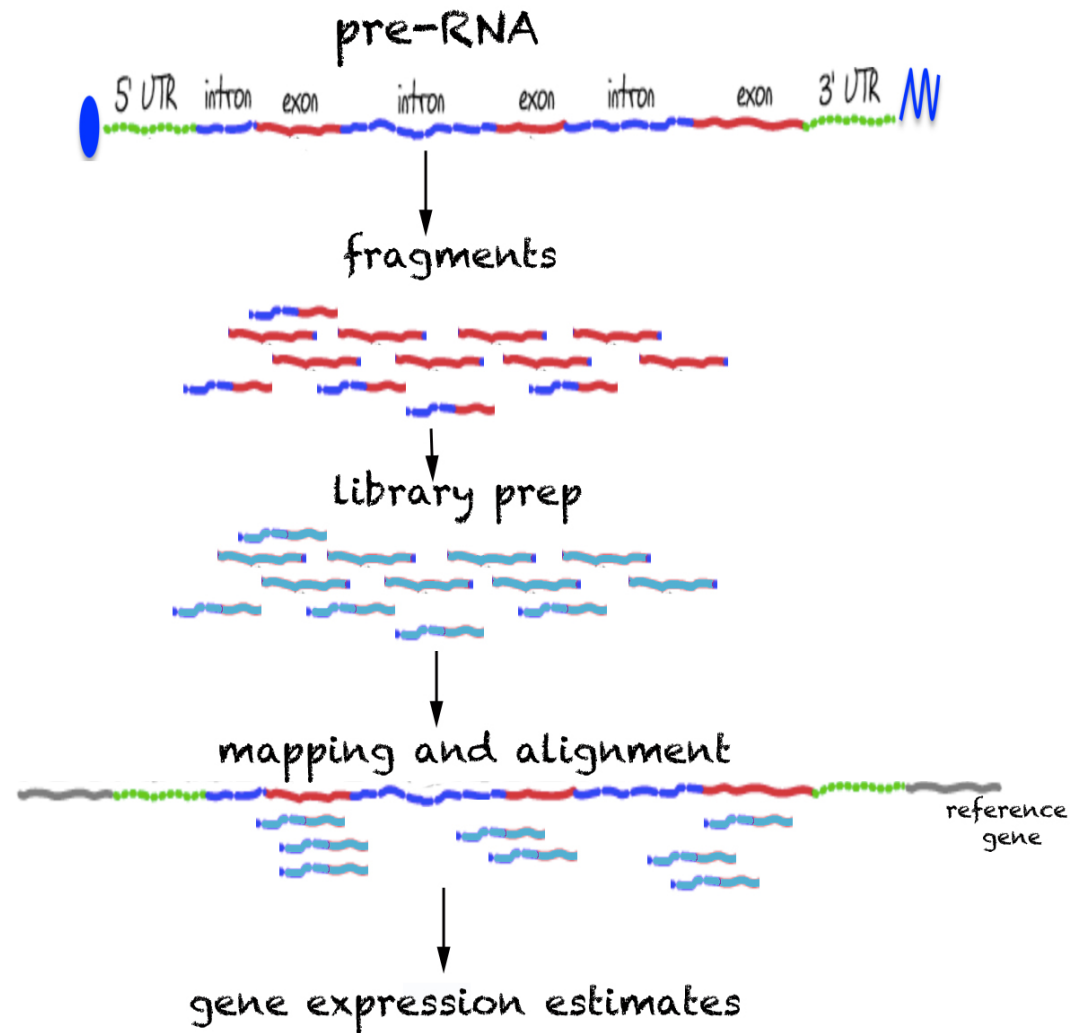
Gene expression quantification using RNA-Seq



High-throughput sequencing of cDNA:

- Shared exons
- Biological variance of fragments
- Splicing variations

Example of RNA-Seq workflow



Illumina sequencing – HiSeq



Illumina HiSeq sequencer

What are we able to detect?

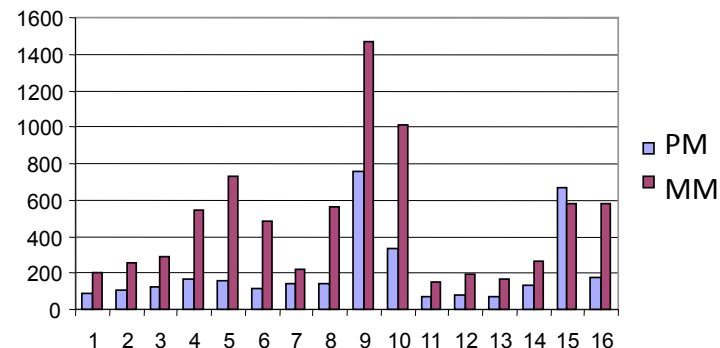
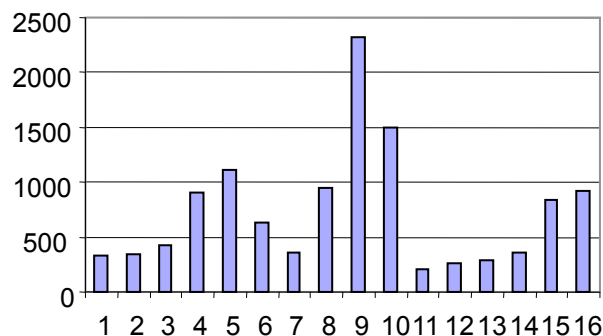
- Mapped read count proportional to abundance of fragments
- Abundance of fragments \approx (gene expression) \times (length)
- Problems which length? which transcripts?
- Other difficulties: mismatches, varying quality of reads, non-uniform read distribution

Part B

What's in the GeneChip data?

1. Summarise to a single expression level the probe intensities for each probe set
2. Estimate the variations introduced by
background effect
probe affinity effect
3. Some PM/MM pairs are more reliable than others
4. The signal needs to be scaled before comparing data from different arrays

How we define a measure that best represent the absolute expression level of each gene on the chip?



The approaches

Use **single point statistics**

make use of the information we have to define values that estimate gene expression

MAS 5.

RMA – GCRMA

PLIER

Use a **probabilistic approach (in Week 10)**

make use of the observed data to estimate function that have generated that data

Estimates of gene expression will be the most probable value that summarises the probe set

PUMA

Data transformation

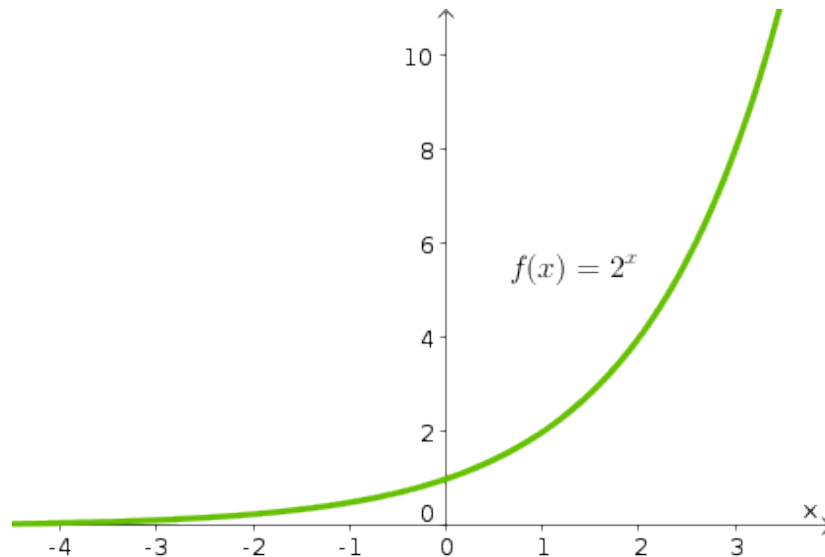
Data transformation converts a set of data values from the format of its original source into a new format that better suits the destination system where the data is then manipulated/used/ mined etc...

In statistics, data transformation refers to the modification of every point in a data set by a mathematical function.

When applying transformations, the measurement scale of the variable is modified.

Data transformation is most often used to change data to the appropriate form for a particular statistical test or method.

Data Transformation: Exponential functions



$$1 + x + x^2 + \dots + x^n$$

$$x^n * x^m = x^{n+m}$$

$$x^n / x^m = x^{n-m}$$

$$y = f(x) = e^x = \exp(x)$$

The number **e** is the Euler's number, an irrational number

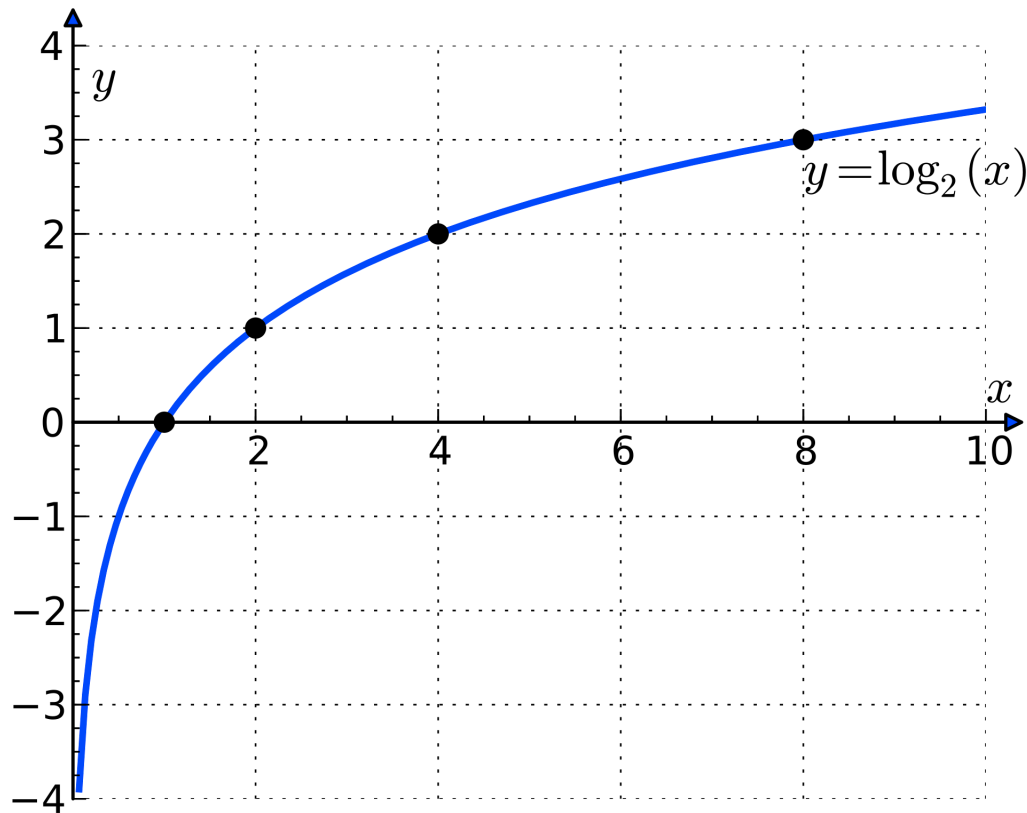
The first few digits are:

2.7182818284590452353602874713527 (and more ...)

Gaussian Function.

$$y = f(x, \mu, \sigma) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x - \mu)^2}{2\sigma^2}\right)$$

Data Transformation: Logarithmic functions



$$\begin{aligned} \log_b(xy) &= \log_b x + \log_b y \\ \log_b(x/y) &= \log_b x - \log_b y \\ \log_b(xn) &= n \log_b x \\ \log_b x &= \log_a x / \log_a b \end{aligned}$$

$y = \log_b x$ if and only if $by = x$,
where $x > 0$, $b > 0$, and $b \neq 1$.

Microarray Suite (MAS5.0)

$$\text{Signal} \sim \text{TukeyBiweight}(\log_2(\text{PM}_j - \text{IM}_j))$$

Correction for global background.- based on 16 sectors on each array

Ideal mismatch (IM) intensity calculated from MM value and subtracted from PM.

- if $\text{MM} < \text{PM}$ then $\text{IM} = \text{MM}$
- if $\text{MM} > \text{PM}$ then $\text{IM} = \text{PM} - \text{correction value}$

- Signal = Smoothed average over PM/MM pairs representing a gene
- Signal is always positive: Absent - Present Call

MAS5: p-value and calls

- First calculate discriminant for each probe pair:
$$R = (PM - MM) / (PM + MM)$$
- Wilcoxon one sided ranked test used to compare R vs tau value and determine p-value
- Present/Marginal/Absent calls are thresholded from p=value above and
 - **Present** $\leq \alpha_1$
 - $\alpha_1 < \mathbf{Marginal} < \alpha_2$
 - $\alpha_2 \leq \mathbf{Absent}$
- Default: $\alpha_1 = 0.04$, $\alpha_2 = 0.06$, $\tau = 0.015$

Not very precise, accurate only when many replicates are available.
Dependent strongly on MM,
Uses linear scaling normalisation

Robust Multi-array Average (RMA)

$$\text{Signal} \sim \text{Tukey}(\log_2(PM_j - \text{bkgd}_j))$$

- Subtract background for each array from PM
- Intensity- dependent normalisation of PM-Bkgd
- Log transform
- Robust multichip analysis of all PM reporters in the set using Tukey median polishing procedure
- Quantile normalisation :Fit all the chips to the same distribution. Scale the chips so that they have the same mean.

Probe Logarithmic Intensity Error (PLIER)

The PLIER algorithm was developed by Affymetrix and released in 2004. It is part of several commercially available

Incorporates experimental observations of feature behavior.

It uses a probe affinity parameter, which represents the strength of a signal produced at a specific concentration for a given probe.

The probe affinities are calculated using data across arrays.

The error model employed by PLIER assumes error is proportional to observed intensity, rather than to background-subtracted intensity.

It assumes that the error of the mismatch probe is the reciprocal of the error of the perfect match probe.

Improved precision over MAS 5

Normalisation

Why do we need to normalise the data?

1. we want to compare across chips
2. we need to ensure that all the data is equally compared across baseline within the chip

Most methods will have normalisation step incorporated, some other will need to perform it after gene expression estimation

Scaling – Mean and Median
Quantile

Loess (not relevant for affymetrix data and sequencing data)

Normalisation: scaling

The assumption that mapping using quantiles or scaling is reasonable is based on the assumption that “most genes don’t change”, and quantiles use this more extensively than scaling.

If this underlying assumption is doubtful, then using the above methods is not advisable.

Simply linearly scale the gene expression so that the overall mean / median are the same.

The median is more scale-invariant, but for the most part there is little practical difference.

Normalisation: quantile

In statistics, *quantile normalization* is a technique for making two distributions identical in statistical properties. When we quantile-normalise a sample distribution to a reference distribution of the same length, we align the sample distribution to the reference so to make them the same.

Assume that the distributions of probe intensities should be completely the same across chips.

Start with n arrays, and p probes, and form a $[p, n]$ matrix X .

Rank first:

Sort the columns of X , so that the entries in a given row correspond to a fixed quantile.

Then align:

Replace all entries in that row with their mean value.

- Visualise the effect: M-A plot

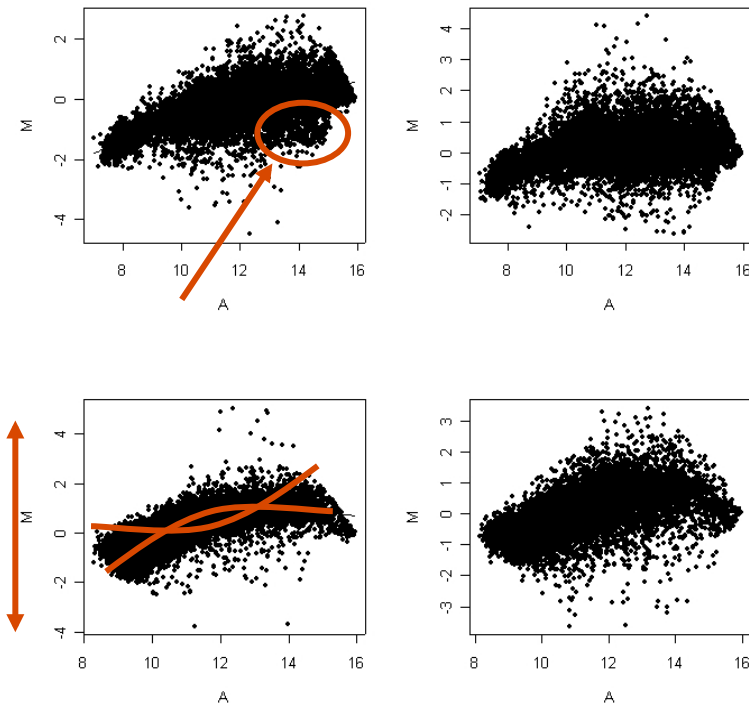
$$M = \log_2 \left(\frac{R}{G} \right)$$

$$A = \log_2 (RG)$$

- Correction of the intensity dependant variations:

$$\log_2 (\text{ratio}) = \log_2 \left(\frac{R}{G} \right) - c(A)$$

$$c(A) = \log_2 k(A)$$



For NGS gene expression quantification....

A lot changes ... but

- Estimation of gene expression
- Alternative splicing identification
- Alternative isoform detection
- Transcripts abundance

QUANTIFICATION



- Normalise read counts
- Normalise reads between lanes
- Normalise reads against transcripts abundance and gene length
- Varying sequencing depth
- Other technical effects

NORMALISATION



Visualise and interpret
Differential expression
Clustering

HIGH LEVEL ANALYSIS

Class activity: debate the following questions:

1. How much do I need to know about the system that I am studying?
2. How much the technologies that are available for data collection need to be sensitive for my system?
3. What is sensitivity and specificity?
4. In sequencing what is a reference genome and how I get it?
5. When the high-throughput approach is the correct approach for my research question?